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# Final Report

## AFOSR Grant F49620-03-1-0385

### Mathematical Modeling of the Continuum Response of Carbon Nanotubes

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### Summary

The authors present a critical analysis of the recent literature related to modeling the continuum mechanical behavior of carbon nanotubes; they also describe a methodology based on combining molecular dynamics simulations, equivalent continua modeling, and mathematical homogenization theory which is directed at obtaining the constitutive relations that can be used to model a nanotube as an elastic cylindrical shell.

## I. Background (The Literature Review)

Interest has been generated in nano-structured materials because of their potential for providing significant improvements in both mechanical and physical properties with respect to traditional structural materials. Investigations [1] point to the fact that carbon nanotubes possess a high elastic modulus (in the 1-5 GPa range) and can sustain an elastic strain on the order of 5% and a fracture strain up to 20%; their strength is, therefore 'unmatched by any other known material' [2]. As indicated in [3] and [4], because small fiber composites are easier to process, the mechanical properties of carbon nanotubes, coupled with their small dimensions and large aspect ratios, make them candidates for the fabrication of the optimal carbon fiber reinforced materials. Work is needed on modeling the constitutive response of single-walled carbon nanotube (SWNT's) and multi-walled carbon nanotubes (MWNT's) in order that reliable constitutive relations can be developed to predict the bulk mechanical properties of nanotube-polymer composites in terms of the molecular structure of the polymer, the nanotubes, and the polymer/nanotube interfaces. As pointed out, in [5], 'in order to implement nanotube-composite design intelligently, it is vital to understand the mechanical behavior of nanotubes.' As the authors of [6] indicate 'even classical molecular dynamics (MD) computations are still limited to simulating on the order of  $10^6 - 10^8$  atoms for a few nanoseconds. The simulation of larger systems or longer times must currently be left to continuum methods.' *The effective continuum modeling of carbon nanotubes, the graphene layers of which they are the 'cylindrical' images, and the delineation of the mapping which takes one from the graphene layer to the nanotube surface, as well as the dependence of that mapping on the chirality vector, form one of the central foci of this proposal, the other being the deduction of a set of constitutive relations for the response of nanotube/polymer composites.*

Almost all modeling of the response of carbon nanotubes ([1], [7]-[25]), and irrespective of whether the authors treat the nanotubes as shells (with some 'effective' thickness) or as beams (with some effective 'cross-section'), *assumes that the 'surface' of the 'cylindrical' nanotube exhibits isotropic mechanical response to deformations*; this assumption is based, in turn, on two hypotheses, namely, that

- (i) the response of a graphene layer is a consequence of the isotropic behavior of graphite ([26], Chapter 3) in the basal plane (or, alternatively, as argued in [9], that the isotropic response of a graphene sheet is a consequence of the intrinsic hexagonal symmetry of the array of carbon atoms comprising the sheet) and
- (ii) that the nanotube can be viewed (e.g., [13]) as a 'conformal mapping of a two dimensional honeycomb lattice to the surface of a cylinder subjected to periodic boundary conditions'

Our research has shown that *neither of hypotheses (i) and (ii) is in accord with the recent literature [6] on MD simulations of the response of carbon nanotubes, or with experimental observations*. Maintaining the isotropy hypothesis for the response of the nanotube in its basal plane has led to *inconsistencies* among the results for the bending stiffness of SWNT's, the representative thickness of nanotubes, and experimental results on in-plane stiffness; these inconsistencies are highlighted in § II. As noted in [6] 'the fundamental assumption of the continuum approximation, that quantities vary slowly over lengths on the order of the atomic scales, breaks down in nano-mechanics.'

Since their discovery there has been an increasing interest in carbon nanotubes because of their novel structure and properties and the wealth of potentially important applications.

As indicated, e.g., in [24], beyond being remarkably stiff and strong, SWNT's are outstanding conductors of electricity and are 'projected to conduct heat even better than diamond.' Under development are SWNT nanodevices and nanocircuits exhibiting superior logic and amplification functions. Because nanotubes are hollow they can be used in sensitive sensor devices as well as in a variety of biomedical applications.

Composite materials based on carbon nanotubes have the potential of providing strength-to-weight ratios which exceed those of any materials currently available, in part because aspect ratios for nanotube fibers of well over 1000 are realizable. As indicated in, e.g., [35], 'in contrast to standard carbon fibers nanotubes remain curved and interwoven in the composite suggesting extreme flexibility.' Effective utilization of nanotubes in composites depends on the ability to disperse the nanotubes homogeneously throughout the matrix (without destroying their integrity). Also, there must be good interfacial bonding between the nanotubes and the polymer to achieve load transfer across the nanotube-matrix interface; the vehicle for such load transfer is the interfacial shear stress. Provided the load can be effectively transferred to the nanotube, the modulus of the composite is conjectured to be that of randomly oriented short fiber composites containing fibers of extremely high modulus and strength. Nanotube composites can be modeled by standard techniques in the theory of heterogeneous materials (e.g., [36] or [37]), or would themselves become a suitable object to which to apply homogenization techniques, *a prerequisite for either approach is a firm understanding of the continuum mechanical behavior of an individual nanotube.* The same considerations apply when one considers MWNT's as potential reinforcing agents; in this case the strength of the composite is also affected by the ease with which individual nanotubes slide with respect to each other. In a MWNT the outer nanotube is loaded but, because of the relatively weak bonding between nanotube layers, the load is not transferred to the inner tubes if they slip with respect to the outertubes; the interactions between SWNT's in a bundle is similar to the interlayer interactions of MWNT's and, as pointed out in [38], slippage within bundles can easily occur. *An effective continuum model for the mechanical response of MWNT's depends on having such a model for SWNT's.*

A graphene sheet is a honeycomb lattice of carbon atoms in a hexagonal array as depicted in fig. 1. In terms of base vectors  $\mathbf{a}_1, \mathbf{a}_2$  the nanotube chirality vector  $\mathbf{r} = n\mathbf{a}_1 + m\mathbf{a}_2$  where  $n, m$  are integers with  $0 \leq |m| < n$ . The chiral angle  $\theta$  is formed by  $\mathbf{r}$  and  $\mathbf{a}_1$ . An  $(n, m)$  nanotube is constructed by using  $\mathbf{r}$  as a 'roll-up' vector, so that with respect to the crystallographically equivalent sites  $A, A', B$ , and  $B'$  ( $\mathbf{t} = \vec{OB}$  is the direction of the nanotube axis)  $A$  and  $A'$  coincide as do points  $B$  and  $B'$ . The rectangle  $AA'B'B$  defines the unit cell for the nanotube. The basic SWNT structures conform to chiral vectors of type  $(n, 0)$  [zig-zag nanotubes] and type  $(n, n)$  [armchair nanotubes]. The rolled-up unit cell is capped by one half of a  $C_{60}$  molecule. Most nanotubes do not exhibit the highly symmetric forms associated with zig-zag and armchair nanotubes but, rather, have structures in which the 'hexagons' are arranged helically around the tube axis (chiral nanotubes). *'Hexagons' lying on the idealized nanotube surface may be deformed versions of those which form the graphitic sheet, e.g., as indicated in [12], with reference to the C-C bonds on the nanotube surface, 'the angles between these bonds depend on the radius of the cylinder as well as on their orientations ... all angles approach 120° (the value in a perfect graphitic plane) with increasing cylindrical radius. For tubules with smaller radii, for example, the bond angles deviate up to 9% from their planar value.'* Moreover, 'upon forming curved surfaces for tubules, the nearest neighbor C-C distances change depending on their relative orientations with respect to the cylindrical axis direction.' Statements in, e.g., [13] which point to the nanotube surface as being 'a conformal mapping

of a two-dimensional honeycomb lattice' *do not accurately depict the deformation which leads from a unit cell of the graphitic sheet to its image on the surface of the (idealized) cylindrical nanotube*. Many authors have suggested using the elastic moduli of graphite for SWNT's by neglecting the change in structure that occurs when a graphene sheet is rolled into a nanotube but recent work [6] has emphasized that '*the deformation is, in fact, not homogeneous as the graphene sheet is rolled into (a carbon nanotube) ... the energy of the (carbon nanotube) not only depends on the deformation gradient but also depends on higher-order derivatives of (the deformation gradient). In such a case, a set of high(er)-order elastic constants that belongs to the framework of multipolar theory needs to be determined.*' Multipolar continuum models have been considered by the authors in [39]-[44]. Thus, *even if we could justify modeling the graphitic sheet as an isotropic elastic continuum one can not carry isotropic response over to the nanotube surface* and recent molecular dynamics simulations (based on Brenner's potential [45]) have been reported by the authors of [6] which lead to a prediction of anisotropic behavior in the basal plane of a nanotube surface.

Other arguments support the hypothesis that the response of the nanotube surface is not that an elastic isotropic material. *If the graphitic sheet can be considered to be an isotropic elastic continuum, whose conformal image is the nanotube surface, then the response of the nanotube should be independent of chirality*: as we roll up a unit cell the graphitic sheet the response of the resulting structure (SWNT) should be independent of the orientation of the edge of the cell with respect to the lattice vector  $a_1$ ; this is not the case, e.g., if one 'rolls up' an arbitrarily oriented rectangular subset of a sheet of rectilinearly orthotropic material, such as paper, into a cylinder (as as been demonstrated by the authors in [46]). Chirality independence has been claimed by several authors, e.g., [9] asserts that *because of the hexagonal symmetry of a graphite sheet its elastic properties are two-dimensionally isotropic, and so the helicity of the tube plays no significant role*" and Lu [8] announced results for carbon nanotubes which are independent of both tube size and chirality. However, it was noted in [19], that '*the apparent insensibility of the Young's modulus on the tube size and chirality observed by Lu is due to the fact that an empirical pair potential was used in his calculations, and such a model will not reflect the effects that the curvature will have on the bonding properties of the system.*' The observations made in [19] have appeared in other recent references, e.g., Yao and Lordi [18] observe that '*for a nanotube of small radius, Young's modulus depends on the radius and helicity of the tube owing to changes in bonding that effect the torsional strain;*' such changes in bonding are directly related to the nonhomogeneity of the map from the graphene sheet to the surface of the nanotube. The failure to include torsional strain contributions in MD simulations of nanotube behavior explains why earlier calculations 'did not reveal significant effects of helicity on elastic properties.' In fact it was noted in [47] that '*the strain energy under large strain conditions shows significant sensitivity to helicity*' while Zhao, et. al. [2] note that '*the behavior of nanotubes under large tensile strain depends strongly on their symmetry and diameter ... different orientations of the carbon bonds with respect to the strain axis in tubes of different symmetry lead to completely different scenarios.*'

One additional set of results mitigate against modeling a SWNT as an elastic shell which is isotropic in its basal plane, namely, *those results which relate Young's modulus, bending stiffness, and in-plane stiffness*. Most authors use the equilibrium interlayer spacing between adjacent nanotubes ( $h = 0.34\text{nm}$ ) as the effective thickness of SWNT's. However, Yakobson, et.al., [9] report an in-plane stiffness for a SWNT of  $C = 59\text{eV/atom} = 360\text{J/m}^2$  and a flexural rigidity of  $D = 0.85\text{eV}$ , as well as a Poisson ration of  $\nu = 0.19$ ; if one uses these measured values *in conjunction with the standard relations associated with an 'isotropic' shell*, i.e.,  $D =$

$Yh^3/12(1-\nu^2)$  and  $C = Yh$  then  $h = 0.066$  nm while the Young's modulus  $Y = 5.5$  GPa. The value of  $h = 0.066$  nm obtained is actually smaller than the typical C-C bond length in the graphitic sheet while the value of  $Y = 5.5$  GPa is several times higher than the usually quoted range of  $1-2$  GPa. However, if (as Ru [24] points out) one begins with a value of  $Y = 1.1$  GPa (consistent with  $h = 0.34$  nm and  $C = 360$  J/m<sup>2</sup>) and substitutes these values into the above expression for  $D$ , a computed  $D$  is obtained which is 25 times larger than the measured value of  $D = 0.85$  eV. These inconsistencies are a consequence of assuming that the nanotube is isotropic in its basal plane. For a cylindrical shell which is the image of a rectilinearly orthotropic sheet one obtains (Bellout and Bloom [46], Bloom and Coffin [48]) for the in-plane principal bending stiffnesses  $D_1 = Yh^3/12(1-(\nu/\gamma)^2)$  and  $D_2 = \gamma^2 Yh^3/12(1-(\nu/\gamma)^2)$  which are equal if and only if the orthotropic ratio  $\gamma = c_{22}/c_{11} = 1$  where  $c_{11}$  and  $c_{22}$  are, respectively, the (in-plane) elastic constants in the axial and radial directions. The construction of a continuum model for the response of a nanotube should not depend on the a priori assumption of basal plane isotropic response and should (i) resolve the inconsistencies with respect to the measured data for  $Y$  and  $C$  and the values of effective shell thickness  $h$  and (ii) provide a basis for the formulation of realistic constitutive relations for nanotube/polymer composites.

## II A Continuum Approach to Nanotube Modelling

In this section we detail an approach to the continuum modeling of single-walled nanotubes; this approach is based on homogenization theory.

Homogenization theory relates to structures that have two naturally associated length scales, a 'micro-scale'  $l$  and and macroscale  $L$ ; the physical properties of the medium vary rapidly on the scale  $l$  and more slowly on the larger scale  $L$ . The ratio  $\epsilon = l/L$  plays a key role wherein it is assumed that every property of the medium is of the form  $\mathbf{F}(\mathbf{x}, \mathbf{y})$  with  $\mathbf{x}$  the position vector of a point in the medium and  $\mathbf{y} = \mathbf{x}/\epsilon$  the vector of stretched coordinates. It is usually the case that the microstructure is periodic (which is reflected in the periodicity of  $\mathbf{F}(\mathbf{x}, \mathbf{y})$  with respect to  $\mathbf{y}$ ); this is the case for the graphitic sheet by virtue of the hexagonal symmetry displayed in fig. 1. For the graphitic sheet  $\mathbf{x} = (x_1, x_2)$ ,  $\mathbf{y} = (x_1/\epsilon, x_2/\epsilon)$  but in its (flat) undeformed state physical properties depend only on  $\mathbf{y}$ , i.e.,  $\mathbf{F} = \mathbf{F}(\mathbf{y})$ .

In homogenization theory each field quantity  $\mathbf{u}^\epsilon(\mathbf{x}) = \mathbf{u}(\mathbf{x}, \mathbf{x}/\epsilon)$  is conceived of as having a double-scale asymptotic expansion of the form

$$\mathbf{u}^\epsilon(\mathbf{x}) = \mathbf{u}^0(\mathbf{x}, \mathbf{y}) + \epsilon \mathbf{u}^1(\mathbf{x}, \mathbf{y}) + \epsilon^2 \mathbf{u}^2(\mathbf{x}, \mathbf{y}) + \dots \quad (1)$$

Each of the terms  $\mathbf{u}^i(\mathbf{x}, \mathbf{y})$  is periodic in  $\mathbf{y}$  with the same period as the structure in question; the ansatz (1) yields, in many cases, a rigorous procedure for deducing macroscopic equations, in  $\mathbf{x}$ , for the overall behavior of the medium.

In the situation we are interested in the global macroscopic behavior is reflected in a set of homogenized elastic constants  $a_{ij}^h$  which mediate the elastic response of a graphitic sheet at the continuum level. At the microscopic level the mechanical response of the graphitic sheet will be governed by a conservation law of the form

$$-\frac{\partial}{\partial x_i} \left( a_{ij}^h(\mathbf{x}) \frac{\partial \mathbf{u}^\epsilon}{\partial x_j} \right) = \mathbf{f}^\epsilon \quad (2)$$

where the  $a_{ij}^\epsilon(\mathbf{x}) = a_{ij}(\mathbf{x}, \frac{\mathbf{x}}{\epsilon})$  are periodic in  $\mathbf{y}$  with period  $\mathbf{Y}$ , i.e.,  $a_{ij}(\mathbf{x}, \mathbf{y}) = a_{ij}(\mathbf{x}, \mathbf{y} + \mathbf{Y})$  and satisfy  $a_{ij}^\epsilon = a_{ji}^\epsilon, a_{ik}^\epsilon \xi_i \xi_k \geq \gamma \xi_j \xi_j, \gamma > 0$ . The period  $\mathbf{Y}$  serves to define a unit cell  $Y$  and will be defined, below, for the graphitic sheet;  $\epsilon$  and  $\mathbf{Y}$  for our problem must be chosen so as to reflect a distinct dependence on the magnitude of the chirality vector as well as on the chiral angle. The (microscale) forcing function  $\mathbf{f}^\epsilon$  is also assumed to be  $\mathbf{Y}$  periodic in  $\mathbf{y}$ .

The basic idea in the homogenization scheme is to substitute the expansion (1) into (2) and then identify powers of  $\epsilon$ ; using the fact that

$$\frac{\partial}{\partial x_j} \phi(\mathbf{x}, \mathbf{x}/\epsilon) = \left( \frac{\partial \phi}{\partial x_j} + \frac{1}{\epsilon} \frac{\partial \phi}{\partial y_j} \right) \quad (3)$$

this procedure yields, at order  $O(\epsilon^{-2})$ ,

$$\frac{\partial}{\partial y_i} \left( a_{ij} \frac{\partial \mathbf{u}^o}{\partial y_j} \right) = \mathbf{0} \quad (4)$$

an equation for the  $\mathbf{Y}$  periodic function  $\mathbf{u}^o(\mathbf{x}, \mathbf{y})$  whose only solution is  $\mathbf{u}^o(\mathbf{x}, \mathbf{y}) \equiv \mathbf{u}^o(\mathbf{x})$ . At order  $O(\epsilon^{-1})$ , assuming  $\mathbf{u}^o(\mathbf{x})$  is known, we obtain for  $\mathbf{u}^1$  the following elliptic equation in a unit cell  $Y$  whose solution must satisfy  $\mathbf{Y}$  periodicity:

$$\frac{\partial}{\partial y_i} \left( a_{ij} \left[ \frac{\partial \mathbf{u}^1}{\partial y_j} + \frac{\partial \mathbf{u}^o}{\partial x_j} \right] \right) = \mathbf{0}. \quad (5)$$

It may be shown (see [27], [28]) that  $\mathbf{u}^1$  can be expressed in the form

$$\mathbf{u}^1(\mathbf{x}, \mathbf{y}) = \frac{\partial \mathbf{u}^o}{\partial x_k} w_k + \hat{\mathbf{u}}^1(\mathbf{x}) \quad (6)$$

where  $\hat{\mathbf{u}}^1(\mathbf{x})$  is an arbitrary function of  $\mathbf{x}$  representing the mean value of  $\mathbf{u}^1(\mathbf{x}, \mathbf{y})$  on  $\mathbf{Y}$  and  $w_k \in W_Y = \{ \phi \in H^1(Y) | \phi \text{ is } Y\text{ periodic and } \hat{\phi} = \frac{1}{|Y|} \int_Y \phi(y) dy = 0 \}$  is the unique solution of the variational equation

$$\int_Y a_{ij} \frac{\partial w_k}{\partial y_j} \frac{\partial \phi}{\partial y_i} dy = - \int_Y a_{ik} \frac{\partial \phi}{\partial y_i} dy, \forall \phi \in W_Y. \quad (7)$$

If we write (2) in divergence form (in terms of the Cauchy stress tensor  $\mathbf{t}$ ) as

$$-\operatorname{div} \mathbf{t}^\epsilon = \mathbf{f}^\epsilon \text{ with } t_{ik}^\epsilon = a_{ij}^\epsilon \frac{\partial u_k^\epsilon}{\partial x_j} \quad (8)$$

expand  $\mathbf{t}^\epsilon$  as  $\mathbf{t}^\epsilon(\mathbf{x}) = \mathbf{t}^o(\mathbf{x}, \mathbf{y}) + \epsilon \mathbf{t}^1(\mathbf{x}, \mathbf{y}) + \dots$ , substitute into the expression for  $\mathbf{t}^\epsilon$  the expansion (1) for  $\mathbf{u}^\epsilon(\mathbf{x})$ , and then set  $\epsilon = 0$ , we obtain

$$t_{ik}^o(\mathbf{x}, \mathbf{y}) = a_{ij} \left( \frac{\partial u_k^o}{\partial x_j} + \frac{\partial u_k^1}{\partial y_j} \right). \quad (9)$$

The integral form of the conservation law in (8), in a macroscopic domain  $D$ , composed of whole periods, is,  $-\int_{\partial D} \mathbf{t}^\epsilon \cdot \mathbf{n} dS = \int_D \mathbf{f}^\epsilon dV$  which, to first order in  $\epsilon$  yields

$$-\int_{\partial D} \hat{\mathbf{t}}^o \cdot \mathbf{n} dS_x = \int_D \hat{\mathbf{f}} d\mathbf{x} \quad (10)$$

From (10) we obtain the *macroscopic equations*

$$-\operatorname{div} \hat{\mathbf{t}}^0 = \hat{f} \text{ with } \hat{t}_{ij}^0 = a_{ik}^h \frac{\partial u_j^0}{\partial x_k} \quad (11)$$

where the *homogenized coefficients (elastic constants)*  $a_{ik}^h$  are given by

$$a_{ik}^h(\mathbf{x}) = \frac{1}{|Y|} \int_Y (a_{ik} + a_{ij} \frac{\partial w_k}{\partial y_i}) d\mathbf{y}. \quad (12)$$

If the original coefficients  $a_{ij} = a_{ij}(\mathbf{y})$  are independent of  $\mathbf{x}$  then the  $a_{ij}^h$ 's are constants; *this will be the case for homogenization of a graphitic sheet because of the strict periodicity of the 'micro' (atomistic) structure*. We observe that the *homogenized medium may be anisotropic even if its behavior at the microstructure level is isotropic*: even if  $a_{ij} = a\delta_{ij}$ , (12) implies that

$$a_{ik}^h = \frac{1}{|Y|} \int_Y a \frac{\partial w_k}{\partial y_i} d\mathbf{y} \quad (13)$$

in which case it *does not necessarily follow that*  $a_{ik}^h = 0$  for  $i \neq k$ . This observation is of central importance for the (homogenized) macroscopic behavior of a graphitic sheet.

*To apply homogenization theory to a 'nanotube unit cell', in an undeformed graphitic sheet, we now identify both the microscale unit cell  $Y$  and the elasticity coefficients  $a_{ij}(\mathbf{x}, \mathbf{y})$  which appear in (12).*

In fig. 2a we depict one 'hexagonal' array of carbon atoms in the unit cell  $AA'B'B$  shown in fig. 1. We have located the vertex at which the bravais lattice vectors  $\mathbf{a}_1, \mathbf{a}_2$  originate at the point  $(0, 0)$  in the  $x, y$  plane. The chiral angle for the array from which this 'hexagon' has been selected is  $\theta$  and the chirality vector  $\mathbf{r}$  has components  $(n, m)$  with respect to the basis vectors  $\mathbf{a}_1, \mathbf{a}_2$ ; the 'hexagon' is shown as having a 'banded' structure with domain  $Y_1$  being an idealized rectilinear domain of width  $\lambda$ . In  $Y_1$  non-zero elastic constants are generated by the forces associated with C-C bonding; these forces are modeled by molecular dynamics simulations. In the 'interior' of the 'hexagon' the elastic constants on the microscale are taken to be zero. We note that (i) *the assumption that the arrays of carbon atoms in the graphitic lattice are perfect 'hexagons' is a mathematical idealization only*, (ii) *the band width  $\lambda$  is not well-defined and may be chosen to be a parameter in the model*, and (iii) *the curves bounding the domain  $Y_1$  need not be rectilinear but may be idealized to be so in a first model*.

To make clear the dependence of the homogenized coefficients  $a_{ij}^h$  on both the chiral angle  $\theta$  and the magnitude  $\sqrt{m^2 + n^2}$  of the chirality vector  $\mathbf{r}$  we proceed as follows: (i) rotate the banded hexagon in fig. 2a.  $\theta$  radians counterclockwise so that the rotated lattice vector  $\mathbf{a}_1$ , i.e.  $\mathbf{a}'_1$  in fig. 2b, is in the direction of the chirality vector  $\mathbf{r}$ , (ii) identify the vector of periodicity as being  $\mathbf{a}'_2$ , select for the microscale measure the length  $l$  of the line segment  $\overline{OA}$  in fig. 2a., (iii) choose  $\epsilon = l/\sqrt{m^2 + n^2}$ , and (iv) define the domain of integration in (13) to be  $Y \equiv Y_\theta = Y_1^\theta \cup Y_2^\theta$  where the  $Y_i^\theta, i = 1, 2$ , as shown in fig. 2b are the rotated images of the planar domains  $Y_i, i = 1, 2$  of fig. 2a. In the present situation (undeformed lattice) the molecular level  $a_{ij}(\mathbf{y})$  are of the form

$$a_{ij}(\mathbf{y}) = \begin{cases} \bar{a}_{ij}(\mathbf{y}), & \mathbf{y} \in Y_1^\theta \\ 0, & \mathbf{y} \in Y_2^\theta \end{cases} \quad (14)$$

While the  $a_{ij}(\mathbf{y})$  may appear to be discontinuous along the boundary between  $Y_1^\theta$  and  $Y_2^\theta$  one

may always introduce standard ‘cut-off’ functions so as to effect a smooth transition from  $Y_1^\theta$  to  $Y_2^\theta$ .

The remaining issue is the specification of the  $\bar{a}_{ij}(\mathbf{y})$  in (15); these are obtained from MD simulations ([49], [50] or [51]) of the energy associated with the C-C bonding in graphitic sheets. The total energy associated with a typical C-C bond has the form

$$E = E_R + E_\theta + E_\emptyset + E_\omega + E_{vdW} + E_{el} \quad (15)$$

where  $E_R$  is associated with bond stretching,  $E_\theta$  with bond angle bending,  $E_\emptyset$  with torsion,  $E_\omega$  with inversions (out-of plane bending),  $E_{vdW}$  with Van der Waals interactions, and  $E_{el}$  with electrostatic interactions. Our work, to this point has singled out functional forms for each term in (15) which are given, e.g., in [38] and are based on MD simulations; alternative forms for  $E$  may be found in [52] which compares the consequences of using either standard ‘Brenner’ or ‘modified Morse’ potentials. Employing the Tersoff-Brenner model the review [6] has shown how to *generate the elasticity tensor associated with any particular MD model by treating the energy E as the potential function for a hyperelastic material; we have now determined that such a procedure will serve well to generate the  $\bar{a}_{ij}(\mathbf{y})$  in (15)*. In particular our early work has directed us to focus on the classical Tersoff-Brenner model with associated empirical equation

$$\Phi_{ij} = \Phi_R(r_{ij}) - \bar{B}_{ij}\Phi_A(r_{ij}) \quad (16)$$

in which  $i, j$  are indices for neighboring carbon atoms,  $r_{ij}$  is the corresponding spatial separation, and  $\Phi_R$  and  $\Phi_A$  represent the repulsive and attractive part of the potential, respectively. The effect of bonding angle is included in the term  $\bar{B}_{ij}$ . The energy density  $E$  is then given by

$$E = \Phi/A_o, \quad \Phi = \sum_{l=1}^3 \Phi_l \quad (17)$$

where  $A_o$  is the area of the undeformed hexagon and  $\Phi_l$  is the bond energy for the  $l$ -th bond. Finally, in terms of the Lagrangian strain tensor  $e_{ij}$ , associated with a deformation of the graphene sheet, we will take for the elasticity tensor components

$$c_{ijkl} = \frac{\partial^2 E}{\partial e_{ij} \partial e_{kl}}$$

which, because of the symmetries with respect to interchange of indices, serves to generate the  $\bar{a}_{ij}$  in (14).

As the graphene sheet is ‘rolled up’ into a nanotube, the hexagonal array in the unit cell  $AA'B'B$  deforms (fig. 3 depicts the different surface arrays for two types of SWNT’s); the deformation gradient associated with this mapping (the mapping  $f : A \rightarrow B$  of fig. 4) is determined, as pointed out in [6], by “the direction and length of the chirality vector and the subsequent mechanical relaxation after rolling.” For various configurations of deformed ‘hexagonal arrays’ on nanotube ‘surfaces,’ one must conformally map those surfaces onto the plane (the mapping  $g : B \rightarrow C$  in fig. 4), apply homogenization to the planar configuration  $C$ , and, subsequently to determine the composition of the mappings  $f$  and  $g$ . The deformation gradient associated with the mapping  $g \circ f$  serves to carry the homogenized coefficients  $a_{ij}^h$ , computed for the graphitic sheet, onto a set of associated homogenized coefficients  $\bar{a}_{ij}^h$  for

the nanotube. While the homogenized coefficients  $a_{ij}^h$  should be constants we expect that  $\tilde{a}_{ij}^h = \tilde{a}_{ij}^h(\mathbf{x})$ . The result of this process may lead to a set of higher-order elastic constants and, thus, to the need for a multipolar theory.

Our work has shown that two classes of results on SWNT shell thickness are prevalent in the literature: *those which assume, a priori, a thickness  $h$  equal to the inter-layer separation of graphitic sheets in a graphite crystal and those which are ‘backed out’ from experimental measurements of in-plane stiffness and flexural rigidity coupled with an assumption of isotropic elastic response.* The in-plane elastic constants in the axial and radial directions for the SWNT,  $c_{11}$  and  $c_{22}$ , are given by  $c_{11} = \tilde{a}_{11}^h$  and  $c_{22} = \tilde{a}_{22}^h$  so that the orthotropic constant  $\gamma = \tilde{a}_{22}^h/\tilde{a}_{11}^h$ ; *this constant can be used to compute the in-plane principal bending stiffnesses  $D_i$  which is used in place of the flexural rigidity associated with isotropic response to ‘back out’ a predicted shell thickness  $h$ .* Such an approach assumes orthotropic response in the basal plane of a SWNT surface (i.e.,  $\tilde{a}_{12}^h = \tilde{a}_{21}^h$ ) as opposed to full anisotropic response. While the principal bending stiffnesses associated with anisotropic response may be computed, *an alternative approach to computing  $h$ , which is based on [31]-[34] may also be used.*

The focus of the work in [31]-[34] is the development of an *equivalent continuum model of a representative volume element (RVE) for a nanotube/polymer composite*; as a first step in the formulation of such a model, an atomistic model of the equilibrium response of the graphitic sheet is obtained by employing MD and expressing the total molecular potential energy in the form (15). A transition is then effected which carries the analysis (see fig. 5) from the MD simulation of the energy for the hexagonal array of carbon atoms (5a), to the energy associated with deformation of an ‘equivalent’ pin-jointed truss (5b), to the strain energy for an ‘equivalent’ hexagonal plate of thickness  $h$  (5c). *Using only the terms,  $E_R$  and  $E_\theta$ , in (15), the energy for the configuration of fig. 5a was taken in the form*

$$E = \sum_a K_a^\rho (\rho_a - P_a)^2 + \sum_a K_a^\theta (\theta_a - \Theta_a)^2 \quad (18)$$

where  $P_a$  and  $\Theta_a$  refer, respectively, to the undeformed interatomic distance of bond number  $a$  and the undeformed bond-angle number  $a$ , while  $\rho_a$  and  $\theta_a$  are, respectively, the interatomic distance and bond angle after deformation. The symbols  $K_a^\rho$  and  $K_a^\theta$  stand for the (MD) force constants associated with stretching and angle variance. *The energy (18) ignores the contribution of torsional strain energy.* Using (18) an equivalent continuum model is developed in [32] in a two-step process; first, the hexagonal array of carbon atoms is replaced by a pin-jointed truss model in which truss elements are used to represent the chemical bonds in the lattice structure. In fig. 5b deformations of rods of type  $a$  are used to describe bond stretching while stretching and contraction of the rods of type  $b$  are used to describe angle variations. The mechanical strain energy for such a truss is

$$\hat{E} = \sum_b \sum_a \frac{A_a^b Y_a^b}{2R_a^b} (r_a^b - R_a^b)^2 \quad (19)$$

where  $A_a^b$  and  $Y_a^b$  are the cross-sectional area and Young’s modulus of rod  $a$  of truss member type  $b$ , respectively, while  $r_a^b - R_a^b$  is the ‘stretching’ of rod  $a$  of truss member type  $b$ , and  $R_a^b, r_a^b$  are the undeformed and deformed lengths, respectively, of the truss elements. *By comparing (18) and (19) the strain energy of the truss model is expressed in terms of the potential energy constants  $K_a^\rho$  and  $K_a^\theta$ .* Finally, the truss model in fig. 5b is replaced by an ‘equivalent’ continuum plate (fig. 5c); the authors *assume the ‘equivalent’ plate to be elastically isotropic*

using for its mechanical properties the Young's modulus and Poisson ratio of graphite. It is also assumed that the same amount of elastic strain energy is stored in the two models (truss model (5b) and continuum plate (5c)) when each is deformed under identical static loading conditions; comparison of the strain energies for the models in figs. 5b,c then yields a result for the thickness of the 'equivalent' continuum plate and, therefore, for the effective thickness of the graphitic sheet. The computations in [32] yield effective thicknesses in the range 0.24nm - 0.28nm, *an improvement over the a priori assumption of 0.34nm but not an improvement which resolves the inconsistencies reported in § II.*

In the work we have initiated (i) *the a priori assumption of a strain energy based on isotropic elastic response for the continuum plate (fig. 5c) is replaced by a strain energy based on the constitutive response predicted by homogenization and (ii) the energy E for the atomistic scale hexagon is expanded to include a suitable MD expression for  $E_\phi$ , the torsional strain energy; equivalent energy terms must be incorporated into the total energy for the pin-jointed truss model and then compared, under similar static loading conditions, with the new elastic strain energy expression for the continuum plate so as to yield a prediction of effective thickness; this will make possible the computation of a realistic Young's modulus.*

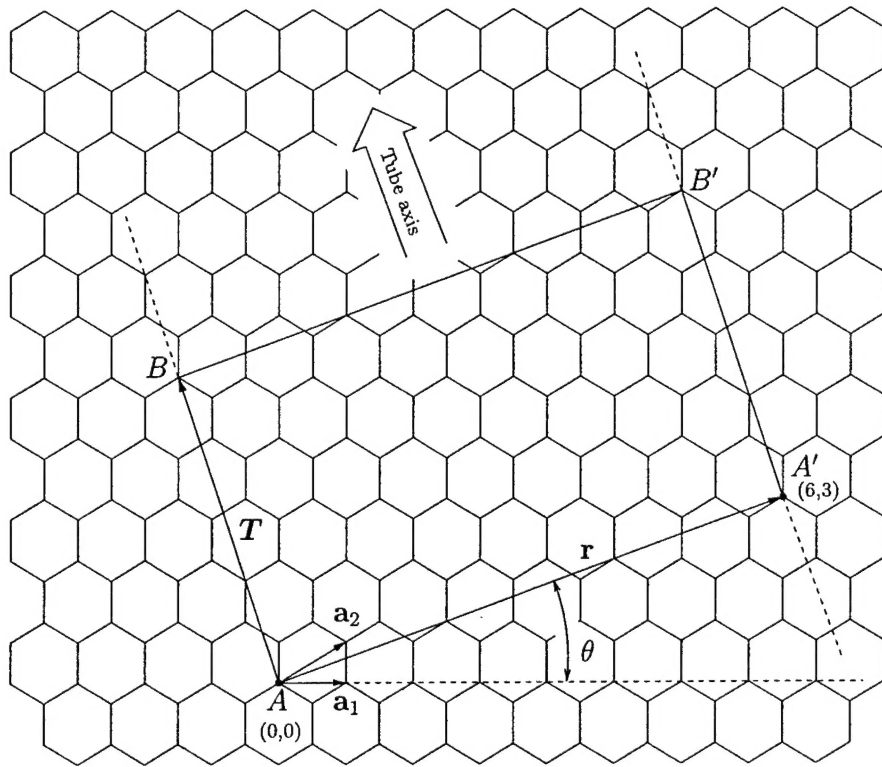
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**Figure 1** (from P.J.F. Harris, Carbon Nanotubes and Related Structures, New Materials for the Twenty-First Century, Cambridge University Press, 2003, Cambridge, U.K.)

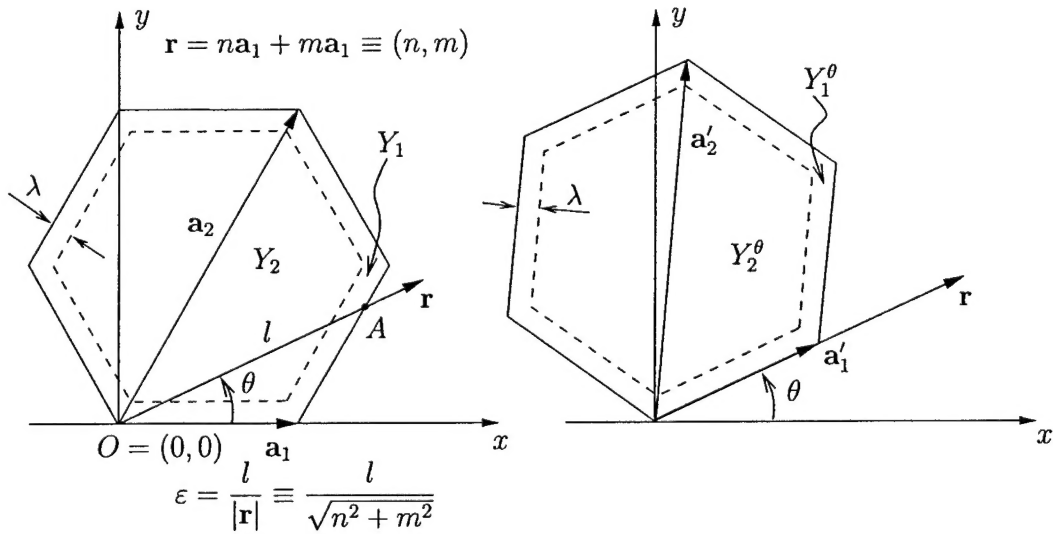


Figure 2a.

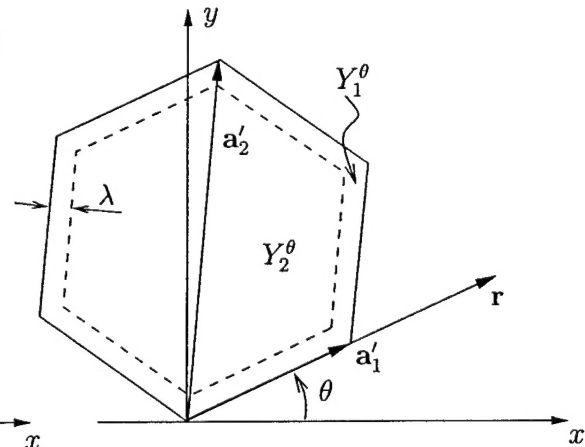
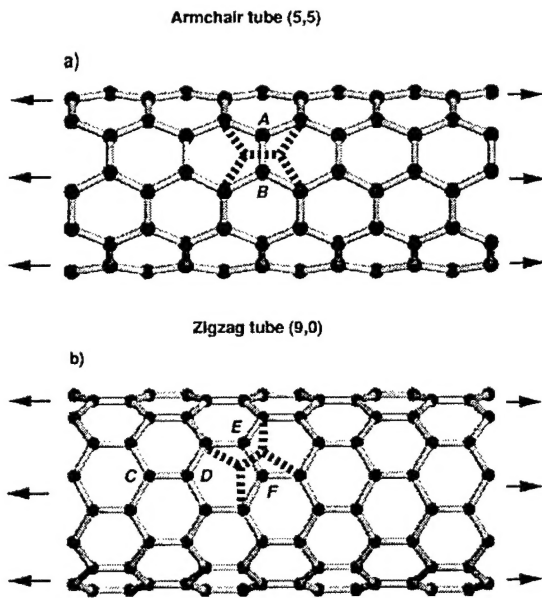
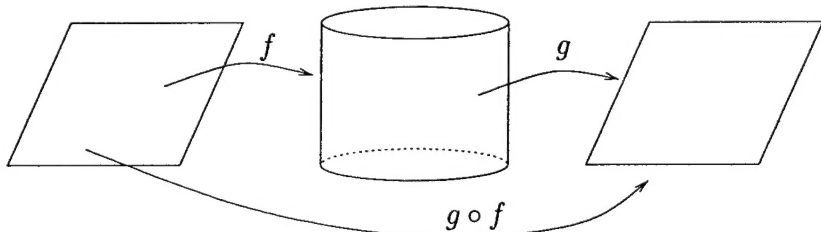


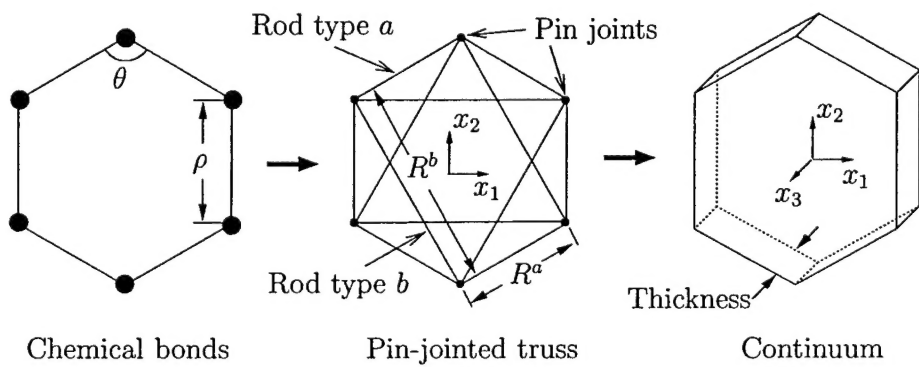
Figure 2b.



**Figure 3.** (from Q. Zhao, M. Nardelli, & J. Bernholc, 'Ultimate Strength of Carbon Nanotubes: A Theoretical Study,' Phys. Rev. B. 65, (2002), 144105-1-144103).



**Figure 4.**



**Figure 5.** (from T.S. Gates & J.A. Hinkley, 'Computational Materials: Modeling and Simulation of Nanostructured Materials and Systems', NASA/TM-2003-212103, NASA Langley Research Center, Hampton, VA)